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## **AMENDMENTS**

## In the abstract:

Methods are provided for identifying heterogeneous features, including heterogeneous background regions, in an image of an array, e.g., in an image of a biopolymeric array, such as a nucleic acid array. The subject methods employ an algorithm that employs a different dispersity measure depending on whether the signal features are weaker or stronger. In the subject methods, a toggle parameter, e.g., a single value (i.e., toggle point) or range of values (i.e., toggle range, smooth function), for the chary of features is first determined. The togglo parameter is determined using statistics obtained fresadow signal features on the array. Following determination of the toggle parameter, there features that have a signal intensity that is either; (a) equal to or less than the toggle parameter and have an intra-feature neise metric-1-level, e.g., standard deviation, that expected the intra-feature noise-limit for metric-1; or (b) greater than the toggle point and have an intra-feature noise metric 2 level-that executs the intra-feature noise limit for metric 2, e.g., ex efficient of variation, we identified as heterogeneous. Also provided are computer readable storage media that include an algorithm capable of performing the steps of the subject recthods. The subject methods find use in the processing of images obtained from a variety of different types of arrays, including nucleic acid arrays.